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㉒ **Masonry treating composition and method.**

㉓ A composition which is an anhydrous and organic solvent free mixture of a hydrolyzable silane monomer ; silanols formed by hydrolysis of the silane monomer ; alkoxy functional oligomers formed by condensation of the silanols such as dimers and trimers and higher alkoxy functional oligomers such as tetramers. Methods of making and using this composition are also described.

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This invention relates to a composition and a method of preparing said composition that are useful in the treatment of porous surfaces such as masonry. More particularly, the composition contains a hydrolyzable silane monomer, silanols and alkoxy functional oligomers and is characterized by being both anhydrous and substantially organic solvent free.

5 The advent of environmental regulations directed to environmental concerns, particularly in the area of architectural coatings, has created the need for solventless compositions. Solvent free architectural coatings containing silanes are known in U.S. Patent No. 4,716,051 issued December 29, 1987. In the '051 patent, 100 percent of a pure solvent free alkyltrialkoxy silane is applied to the surface of concrete to render the surface hydrophobic.

10 The present invention is an improvement compared to this technology and provides a treating composition which is anhydrous as well as being solventless or free of organic solvent, while at the same time including in addition to a silane monomer, silanols and alkoxy functional oligomers. The prehydrolyzed composition of the present invention will generate less organic alcohol liberated during further hydrolysis of alkoxy groups upon contact with moisture, thereby rendering the composition of the present invention more environmentally acceptable. Since the compositions are also anhydrous, stability problems during storage of the composition and 15 prior to its use, which are caused by decomposition in aqueous medium are reduced. Thus, the compositions will remain stable for a sufficient time to make it economically feasible to employ these materials in a variety of coating applications.

15 The invention is directed to a composition for the treatment of porous surfaces such as masonry which is an anhydrous organic solvent free mixture containing (i) a hydrolyzable silane monomer; (ii) silanols of the silane monomer formed by partial hydrolysis thereof; (iii) lower alkoxy functional oligomers formed by partial condensation of the silanols including dimers and trimers thereof; and (iv) higher alkoxy functional oligomers formed by partial condensation of the silanols including tetramers thereof.

20 The invention is further directed to a method of preparing this composition. The method involves the steps of forming a solution by combining water with a hydrolyzable silane monomer in a range of 0.1 to 0.6 moles of water per mole of silane monomer in the presence of an acid catalyst and partially hydrolyzing the silane monomer to form silanols and alcohol while simultaneously partially condensing the silanols to form alkoxy functional oligomers; neutralizing the solution by adding a mild base to the solution; heating the solution to a temperature sufficient to strip the alcohol from the solution and separating any excess mild base and any neutralization 25 products from the solution. The product isolated is an anhydrous organic solvent free mixture of residual hydrolyzable silane monomer, residual silanols and lower alkoxy functional oligomers including dimers, trimers and higher alkoxy functional oligomers including tetramers.

30 The invention in addition is directed to a method of treating a porous surface such as masonry for the purpose of rendering it hydrophobic by applying to the porous surface a composition which is an anhydrous organic solvent free mixture of a hydrolyzable silane monomer, silanols and lower and higher alkoxy functional oligomers such as dimers, trimers, tetramers.

35 The problem solved by the present invention is to provide a method of making a masonry treating composition in which a critical range of moles of water per mole of silane monomer results in the provision of an end product, which is anhydrous and organic solvent free and in compliance with environmental standards of 40 current architectural coating regulations.

45 Typically, the hydrolysis of the alkoxy groups of an organotrialkoxy silane requires molecular contact of water with the alkoxy silane in the presence of an acid or a base catalyst. Organotrialkoxysilanes are commonly dissolved in water by shaking or stirring vigorously with acidified water until a clear solution results. Trialkoxysilanes of the formula $RSi(OR')_3$ hydrolyze stepwise in water to the corresponding silanols which condense to siloxanes. The hydrolysis reaction is relatively rapid while the condensation reaction is slower.

50 The former reaction is typically conducted in the presence of an acid catalyst which accelerates the hydrolysis. Preferred acid catalysts are hydrochloric, acetic, oxalic and trichloroacetic acids, with hydrochloric acid being the most preferred. A strong basic catalyst may also be employed. Representative strong bases are sodium hydroxide and sodium methylate.

55 For purposes of the present invention, the hydrolyzable silane monomer $RSi(OR')_3$ is an organotrialkoxy silane or a mixture of organotrialkoxysilanes such as alkyltrialkoxy silanes having C_1 to C_{20} alkyl groups on silicon; aryltrialkoxy silanes; arylalkyltrialkoxy silanes and halogen substituted derivatives thereof. Representative compounds which may be employed in accordance with the present invention are methyltrimethoxysilane, ethyltrimethoxysilane, propyltrimethoxysilane, isopropyltrimethoxysilane, butyltrimethoxysilane, isobutyltrimethoxysilane, pentyltrimethoxysilane, hexyltrimethoxysilane, phenyltrimethoxysilane, phenylethyltrimethoxysilane, chlorophenyltrimethoxysilane, trifluoropropyltrimethoxysilane and chloropropyltrimethoxysilane.

60 In addition to the foregoing $RSi(OR')_3$, there may also be employed silanes in which R' is represented by an alkyl group having one to eight carbon atoms, preferably one to four carbon atoms or phenyl. Thus, approx-

5 appropriate hydrolyzable silane monomers for purposes of the present invention may include the various ethoxy, isopropoxy and phenoxy derivatives of the several silanes enumerated previously. In addition, mixtures of the silanes may be employed where it is desired to produce co-oligomeric products. In this event, the mixture can for example include a combination of an alkyltrialkoxysilane such as isobutyltrimethoxysilane, with a halogen substituted derivative such as trifluoropropyltrimethoxysilane. While this is the preferred combination, it should be understood that other various mixture combinations may be utilized.

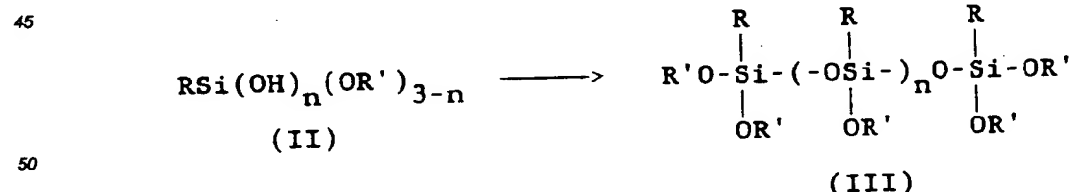
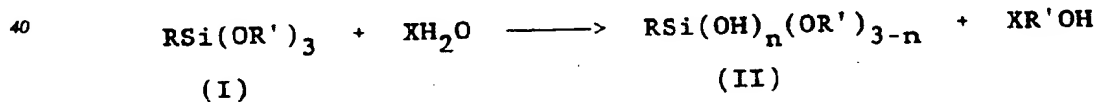
10 Following hydrolysis and condensation, the solution must be freed of hydrochloric acid and this is accomplished by neutralization with a mild base such as calcium carbonate in contrast to strong bases such as sodium hydroxide and sodium methylate, although other neutralizing agents classified as mild bases may be employed such as sodium carbonate, sodium bicarbonate, ammonium carbonate, sodium borate, sodium acetate, calcium acetate, calcium hydroxide, potassium acetate and ammonium acetate. A mild base is required because over-neutralization with a strong base can create a product which is unstable. Following neutralization, the solution is distilled to remove the byproduct alcohol formed during hydrolysis and the distilled solution is filtered in order to separate the excess neutralizing agent and any neutralization products from the final composition.

15 A critical feature of the present invention resides in the particular amount of water employed for the hydrolysis in relation to the amount of silane monomer. It has been found that compositions in accordance with the present invention may be produced by employing a very specific and limited range of moles of water per mole of silane, which range may vary between 0.1 to 0.6 moles of water per mole of silane. The range preferred is 0.25 to 0.5 moles of water per mole of silane, with the most preferred amount being 0.5 moles of water per mole of silane. Lower amounts of water favor the formation of the lower alkoxy functional oligomeric dimers and trimers, while high amounts of water provide higher alkoxy functional oligomers such as the tetramers, pentamers and higher species.

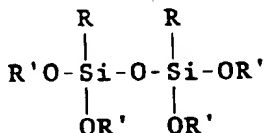
20 Thus, a process suitable for the preparation of compositions in accordance with the present invention is carried out by the steps of forming a solution by combining a limited amount of water to a neat hydrolyzable silane monomer in the presence of an acid catalyst; stirring the solution; neutralizing the solution; stripping byproduct alcohol from the solution by heating the solution at atmospheric pressure to a temperature of about 150°C. and separating from the solution any solid byproducts. Condensation of the hydroxyl radicals formed by hydrolysis of the silane monomer may be facilitated by applying external heat to the aqueous reaction solution for a period of time prior to neutralization.

25 30 Where a limited amount of water in the range of 0.1 to 0.6 moles of water per mole of silane is employed in the process as outlined above, the final composition produced will constitute an anhydrous organic solvent free mixture which includes 10.0 to 65.0 percent by weight of residual hydrolyzable silane monomer, more than zero but less than about 0.1 percent by weight of residual silanol; 25.0 to 80.0 percent by weight of lower alkoxy functional oligomers including dimers and trimers and 0.25 to about 10.0 percent by weight of higher alkoxy functional oligomers including tetramers.

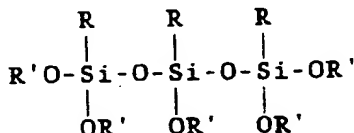
35 By employing a limited amount of water, the scenario described hereinabove is altered and the hydrolysis and condensation reaction results in products being formed in accordance with the following reaction scheme:



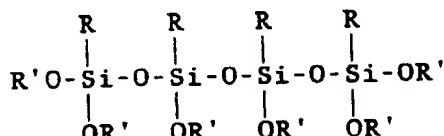
50 The lower and higher alkoxy functional dimer, trimer and tetramer, are shown below:



Dimer



15 Trimer



25 Tetramer

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The compositions of the present invention may be employed in the treatment of various porous surfaces. 30 While intended primarily for the treatment of concrete and masonry products, other porous surfaces which may be treated encompass textiles; paper; paperboard; leather products and cellulosic materials.

Concrete and masonry surfaces which may be treated include heavy and light weight concrete; gypsum; concrete blocks; soft mud bricks; sand lime bricks; drain tiles; ceramic tiles; sandstone; plaster; clay bricks; natural stones and rocks; roofing tiles; calcium silicate bricks; asbestos cement; slag stones and bricks; stucco; limestone; macadam; marble; grout; mortar; terrazzo; clinker; pumice; terra cotta; porcelain; adobe; coral; dolomite and asphalt. Noncermentitious surfaces which are porous are included such as perlite; cellular glass; vermiculite; mica and diatomaceous earth.

Examples of leather products suitable for treatment are garments, shoes and boots. Textiles include awnings, tents, tarpaulins, rainwear, covers, slickers, canvas, asbestos, fiberglass, natural fibers, peat moss, natural and synthetic yarns, woven and nonwoven materials, carpets and carpet fibers. Porous cellulosic surfaces suitable for treatment with the compositions of the present invention are wood, wood products, fiberboard, cedar, redwood, fir, plywood and structural timbers.

The compositions in accordance with the present invention liberate only limited amounts of volatile materials and hence comply with various regulations regarding "volatile organic content" and "volatile organic compounds". Such regulations typically prohibit a volatile organic content for an architectural coating for example in excess of about four hundred grams per liter. In contrast, many coatings of the prior art containing solvent based alkoxysilanes liberate significant amounts of an alcohol which is a volatile organic compound. The volatile organic content of such prior art solvent-based coatings can be of the order of magnitude of 650-700 grams per liter. Air pollution regulations limit the amount of organic solvents that can be discharged into the atmosphere. The term used for solvents is "volatile organic compounds" (VOC). A volatile organic compound (VOC) is defined as any compound of carbon that has a vapor pressure greater than 0.1 millimeter of mercury (0.01 kPa) at a temperature of 20°C. and a pressure of 760 millimeters mercury (101.1 kPa).

"Volatile organic content" has been defined as the amount of volatile organic compounds (VOC) liberated from a coating as determined by ASTM D3690 and EPA Reference Method 24 which are standard industrial tests. Under the definition, a volatile organic compound is any compound which enters the atmosphere and photochemically reacts in the atmosphere with nitrogen oxides to reduce ozone and form photochemical smog.

Reduction of VOC has been mandated in several jurisdictions and one such regulation requires less than about four hundred grams of volatiles per liter of product to enter the atmosphere. This can be determined by

baking ten grams of a product in an oven at 110°C. for one hour. The amount of solids which remain is subtracted from the total of the ten grams which was tested. Calculations are based on the weight of the volatiles that have evaporated which is reported as grams per liter.

The U.S. Environmental Protection Agency (EPA) has identified over sixty different volatile organic compounds that are used in consumer products. The more common solvents found are ethanol, isopropyl alcohol, kerosene and propylene glycol, as well as hydrocarbon solvents such as isobutane, butane and propane which are also employed as propellants.

Some governments have proposed standards which would limit and reduce the amount of volatile organic compounds (VOC) permitted in various consumer products, such as chemically formulated products used by household and institutional consumers including detergents; cleaning compounds; polishes; floor products; cosmetics; personal care products; home, lawn and garden products; disinfectants; sanitizers and automotive specialty products. These standards would affect such widely used consumer products as shaving lather, hair-spray, shampoos, colognes, perfumes, aftershave, deodorants, antiperspirants, suntan preparations, lotions, breath fresheners and room deodorants. Thus, the need for new and novel formulations and techniques for reducing organic emissions should be more than apparent.

The porous surface to be treated by the composition of this invention can be coated by brushing, dripping or spraying techniques, the coating being thereafter allowed to migrate within the surface. Further hydrolysis and condensation may occur because of moisture in the atmosphere and moisture contained in the porous surface itself or because of the highly alkaline nature of the surface being treated. When applied to a porous surface such as masonry, the alkoxy functional oligomers (III) are capable of reacting with OH groups present within the surface. The presence in the composition of residual silanol (II) will tend to increase the hydrogen bonding and provide the benefit of lowering the evaporation of the composition from the substrate. The silanols (II) will also enable the composition to become bound to the substrate more rapidly.

The following examples are set forth in order to further illustrate the concepts embodied in the present invention.

EXAMPLE I

Into a one thousand milliliter flask there was added 2.6 gram moles of isobutyltrimethoxysilane (IBTMS). The contents of the flask were agitated and there was added to the flask 3.5 grams of a 36 percent hydrochloric acid solution for the purpose of catalyzing the hydrolysis. Water was slowly added to the flask at ambient temperature (25°C.) in the amount of 1.2 gram moles. As shown in Table I, three water ratios were evaluated as indicated by Samples A-D. Sample "B" it is noted contained 1.5 times the amount of hydrochloric acid as Samples "A", "C" and "D". The reaction mixture exothermed during the addition of water to the flask and the maximum temperature attained was 39°C. The mixture was allowed to equilibrate at ambient temperature for four hours. The composition of the hydrolysis products for the four samples A-D are shown in Table I. As previously noted, the preferred method for neutralizing the acid catalyst following hydrolysis is by the use of a mild base such as calcium carbonate. Accordingly, 7.0 grams of calcium carbonate were added to the agitated flask. For purposes of comparison, 3.8 grams of sodium methylate (CH_3ONa) a strong base, were used to neutralize the catalyst in Sample "G" as shown in Table I. Sodium methylate caused deviations from the neutral point and produced an unstable product during the distillation phase. Seven grams of activated charcoal were used for the removal of chromophors from the product. The mixture was allowed to equilibrate at ambient temperature for four hours. The mixture in the flask was slowly heated at atmospheric pressure for the purpose of removing methanol generated during hydrolysis. The methanol was collected in another flask following condensation in a tap water cooled condenser. Distillation was discontinued when the liquid in the flask had reached 160°C. The flask was allowed to cool to ambient temperature. The product in the flask was filtered for the purpose of removing calcium carbonate, activated charcoal and calcium chloride salts formed in the neutralization step of the process. Filtration was accomplished using a laboratory scale pressure filter containing a one micrometer rated filter paper. The filtrate was a clear water white liquid having the composition shown in Table I Samples E-G.

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TABLE I

Material	Moles H ₂ O Per Mole IBTMS	Methanol Percent	Monomer Percent	Dimer Percent	Trimer Percent	Tetramer Percent	Higher Oligomers Percent	Neutralization Method	Residue Percent	
									Percent	Method
IBTMS Feedstock	---	1.19	97.37	1.14	0.3	---	---	---	---	---
IBTMS Hydrolysis Product									0.2	---
A	0.5	21.1	15.33	41.72	16.49	3.94	0.2	2.08	2.08	---
B	0.5	26.46	9.06	37.63	19.6	5.17	---	---	---	---
C	0.375	14.51	33.1	38.28	6.79	0.56	---	---	---	---
D	0.25	9.61	59.1	20.66	2.59	0.24	---	---	---	---
Final Product									2-3	2-3
E	0.5	---	14.0	56.4	22.73	5.58	3.29	CaCO ₃	2-3	2-3
F	0.55	---	9.87	51.2	28.33	8.16	2.44	CaCO ₃	12	12
G	0.5	1.2	30.34	26.12	19.8	13.3	9.32	NaOMe		

The amount of water employed in the process is a critical feature of the present invention. In order to obtain

Note 1: Compositions determined by TCD Gas Chromatograph. Confirmed by GC mass spectroscopy.

Note 2: Silanol content of hydrolysis product free of methanol = 1000-1500 ppm (0.1%).

Note 3: IBTMS = isobutyltrimethoxysilane.

Note 4: NaOMe = sodium methylate.

Note 5: Residue = pentamers, hexamers, etc.

the most desirable distribution of alkoxy functional oligomers in the product, water should be employed in an amount of 0.1-0.6 moles of water per mole of silane monomer. Preferably the amount of water should be 0.25-0.5 moles of water per mole of silane monomer and most preferably 0.5 moles of water per mole of silane monomer. Samples A-D in Table I reflect the alkoxy functional oligomer distributions obtained with each of three ratios falling within the foregoing criteria.

5 Samples "A" and "B" in Table I differ from one another in that the amount of hydrochloric acid employed to catalyze the hydrolysis was 1.5 times more in "B" than in "A". It was noted that higher levels of chloride ion in the system increased the rate of hydrolysis which in turn affected the oligomer distribution. Higher levels of hydrochloric acid favor the formation of a higher alkoxy functional oligomer containing distribution. Thus, Sample 10 "B" can be seen to reflect a lower level of monomer and a higher level of alkoxy functional oligomers following hydrolysis than Sample "A".

15 The use of a mild base such as calcium carbonate in order to neutralize the hydrolysis product is a critical feature in the maintenance of a stable product. The use of a mild base as in Samples "E" and "F" in Table I enables a molar excess of the base to be present during the distillation phase without a continuation of the reaction. However, a strong base such as sodium methylate in Sample "G" in Table I demonstrated that catalysis of the hydrolysis continued in the alkaline system producing higher alkoxy functional oligomers. This is reflected in the "Residue" analysis in Table I showing 3-4 times the amount of non-volatile residue for Sample 20 "G" in contrast to Samples "E" and "F". If not completely removed prior to distillation, strong bases such as sodium hydroxide and sodium methylate foster the formation of higher molecular weight species, rendering it difficult to obtain a stable product. In contrast, a mild base enables one to more carefully stabilize the distribution of oligomers in the final product.

25 The amount of the acid catalyst employed in accordance with the process of the present invention ranges from 250 to 1,000 parts per million. Neutralization is effected with the mild base in an amount of nearly 1.5 times the molar equivalent required.

30 The following example shows that by starting with a mixture of monomeric alkoxy silanes, alkoxy functional co-oligomers can be formed in accordance with the hydrolysis procedure in Example I. In Example II, trifluoropropyltrimethoxysilane referred to in Table II as $F_3PrSi(OMe)_3$, was mixed with isobutyltrimethoxysilane (IBTMS). The progress of the reaction was indicated by monitoring the level of monomer species and the formation of alkoxy functional oligomer combinations following hydrolysis. Table II shows the composition of the two feedstocks, the alkoxy functional co-oligomer hydrolysis product and the final alkoxy functional co-oligo- 35 mer product.

EXAMPLE II

35 Into a one thousand milliliter flask there was added 22.6 gram moles of isobutyltrimethoxysilane (IBTMS) and 0.92 gram moles of trifluoropropyltrimethoxysilane $F_3PrSi(OMe)_3$. The contents of the flask were agitated and there was added to the flask 25.0 grams of a 36 percent hydrochloric acid solution for the purpose of catalyzing the hydrolysis. Water was slowly added to the flask at ambient temperature (25°C.) in the amount of 11.1 gram moles. The reaction mixture exothermed during the addition of water to the flask and the maximum 40 temperature attained was 39°C. The mixture was allowed to equilibrate at ambient temperature for four hours. Thirty-five grams of calcium carbonate was added to the agitated flask, accompanied by the addition of 30.0 grams of activated charcoal for the removal of chromophors from the product. The mixture was allowed to equilibrate at ambient temperature for 4 hours. The mixture in the flask was slowly heated at atmospheric pressure for the purpose of removing methanol generated during hydrolysis. The methanol was collected in another 45 flask following condensation in a tap water cooled condenser. Distillation was discontinued when the liquid in the flask had reached 160°C. The flask was allowed to cool to ambient temperature. The product in the flask was filtered for the purpose of removing calcium carbonate, activated charcoal and calcium chloride salts formed in the neutralization step of the process. Filtration was accomplished using a laboratory scale pressure filter containing a one micrometer rated filter paper. The filtrate was a clear water white liquid having the composition shown in Table II. Silanol content of the product was 1400 parts per million.

TABLE II

Material	Methanol	IBTMS Monomer %	F ₃ PrSi(OMe) ₃ Mono- mer %	Higher Oligomers %
IBTMS Feedstock	1.19	97.37	----	1.4
F ₃ PrSi(OMe) ₃ Feedstock	0.63	----	94.6	4.77
Co-oligomer Hy- drolysis Product	15.0	10.2	0.25	74.55
Co-oligomer Fi- nal Product	----	11.2	----	88.78

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Claims

1. A method of making a mixture containing an alkoxy silane, alkoxy functional oligomers and silanols comprising the steps of forming a solution by combining water with a hydrolyzable silane monomer in a proportion of 0.1 to 0.6 moles of water per mole of silane monomer; adding a catalyst to the solution and partially hydrolyzing the silane monomer to form silanols and alcohol; simultaneously partially condensing the silanols to form alkoxy functional oligomers; neutralizing the solution by adding a mild base to the solution; heating the solution to a temperature sufficient to remove the alcohol from the solution; separating the mild base and any products of neutralization from the solution and isolating a product in the form of an anhydrous organic solvent free mixture of residual hydrolyzable silane monomer, silanols formed by hydrolysis of the silane monomer, lower alkoxy functional oligomers formed by condensation of the silanols including dimers and trimers and higher alkoxy functional oligomers formed by condensation of the silanols including tetramers and higher species.
2. The method according to claim 1 in which the hydrolyzable silane monomer is an organotrialkoxysilane or a mixture of organotrialkoxysilanes selected from alkyltrialkoxysilanes having C₁ to C₂₀ alkyl groups on silicon; aryltrialkoxysilanes; arylalkyltrialkoxysilanes and halogen substituted alkyl and aryl derivatives thereof.
3. The method according to claim 2 in which the organotrialkoxysilane is a compound selected from methyltrimethoxysilane, ethyltrimethoxysilane, propyltrimethoxysilane, isopropyltrimethoxysilane, butyltrimethoxysilane, isobutyltrimethoxysilane, pentyltrimethoxysilane, hexyltrimethoxysilane, phenyltrimethoxysilane, phenylethyltrimethoxysilane, chlorophenyltrimethoxysilane, trifluoropropyltrimethoxysilane, chloropropyltrimethoxysilane, the alkoxy equivalents thereof having from two to eight carbon atoms and aryloxy equivalents.
4. The method according to claim 1 in which the product includes 10.0 to 65.0 percent by weight of residual hydrolyzable silane monomer; more than zero but less than 0.1 percent by weight of silanols; 25.0 to 80.0 percent by weight of lower alkoxy functional oligomers including dimers and trimers and 0.25 to about 10.0 percent by weight of higher alkoxy functional oligomers including tetramers and higher species.
5. The method according to claim 1 in which the catalyst is selected from hydrochloric acid, acetic acid, oxalic acid and trichloroacetic acid and the mild base is selected from calcium carbonate, sodium carbonate, sodium bicarbonate, ammonium carbonate, sodium borate, sodium acetate, calcium hydroxide, calcium acetate, potassium acetate and ammonium acetate.
6. The method according to claim 1 in which the hydrolyzable silane monomer, the silanols, the alkoxy functional oligomers including the dimers, trimers, tetramers and higher species, are methoxy or ethoxy functional.
7. The method according to claim 5 in which the catalyst is hydrochloric acid; the mild base is calcium carbonate and the mild base and any neutralization products are separated from the product by filtration.

8. The method according to claim 7 in which the hydrolyzable silane monomer is contacted with water in a proportion of 0.25 to 0.5 moles of water per mole of silane monomer.

5 9. A method for making a masonry treating composition which comprises contacting a hydrolyzable silane monomer with water in a proportion of 0.1 to 0.6 moles of water per mole of silane monomer in the presence of a catalyst to hydrolyze the silane monomer and condense the hydrolysis product obtained; neutralizing the product obtained by the addition of a mild base to the product; heating the product to a temperature sufficient to remove any alcohol present in the product; separating the mild base and any products of neutralization from the product and isolating the product in the form of an anhydrous organic solvent free mixture.

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